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or a declared completed PDP in effect before being placed in commercial distribution.

[52 FR 33702, Sept. 4, 1987, as amended at 61 FR 50711, Sept. 27, 1996]

§888.3650 Shoulder joint metal/polymer non-constrained cemented prosthesis.

- (a) Identification. A shoulder joint metal/polymer non-constrained mented prosthesis is a device intended to be implanted to replace a shoulder joint. The device limits minimally (less than normal anatomic constraints) translation in one or more planes. It has no linkage across-the-joint. This generic type of device includes prostheses that have a humeral component made of alloys, such as cobalt-chromium-molybdenum, and a glenoid resurfacing component made of ultrahigh molecular weight polyethylene, and is limited to those prostheses intended for use with bone cement (§ 888, 3027)
- (b) Classification. Class II. The special controls for this device are:
 - (1) FDA's:
- (i) "Use of International Standard ISO 10993 'Biological Evaluation of Medical Devices-Part I: Evaluation and Testing,
- (ii) "510(k) Sterility Review Guidance of 2/12/90 (K90-1),
- (iii) "Guidance Document for Testing Orthopedic Implants with Modified Metallic Surfaces Apposing Bone or Bone Cement,
- (iv) "Guidance Document for the Preparation of Premarket Notification (510(k)) Application for Orthopedic Devices," and
 (v) "Guidance Document for Testing
- Non-articulating, 'Mechanically Locked' Modular Implant Components,
- (2) International Organization for Standardization's (ISO):
- (i) ISO 5832-3:1996 "Implants for Surgery—Metallic Materials—Part Wrought Titanium 6-Aluminum Vandium Alloy,
- (ii) ISO 5832-4:1996 "Implants for Surgery—Metallic Materials—Part 4: Cobalt-Chromium-Molybdenum Casting Alloy,
- (iii) ISO 5832-12:1996 "Implants for Surgery—Metallic Materials—Part 12:

Wrought Cobalt-Chromium-Molybdenum Alloy,'

- (iv) ISO 5833:1992 "Implants for Surgery—Acrylic Resin Cements,'
- (v) ISO 5834-2:1998 "Implants for Surgery—Ultra-high Molecular Weight Moulded Polyethylene—Part Forms.
- (vi) ISO 6018:1987 "Orthopaedic Implants—General Requirements Marking, Packaging, and Labeling,'
- (vii) ISO 9001:1994 "Quality Systems-Model for Quality Assurance in Design/ Development, Production, Installation, and Servicing," and ' and
- (3) American Society for Testing and Materials':
- (i) F 75-92 "Specification for Cast Cobalt-28 Chromium-6 Molybdenum Alloy for Surgical Implant Material,'
- (ii) F 648-98 "Specification for Ultra-High-Molecular-Weight Polyethylene Powder and Fabricated Form for Surgical Implants,
- (iii) F 799–96 "Specification for Cobalt-28 Chromium-6 Molybdenum Alloy Forgings for Surgical Implants,"
 (iv) F 1044-95 "Test Method for Shear
- Testing of Porous Metal Coatings,
- (v) F 1108-97 "Titanium-6 Aluminum-4 Vanadium Alloy Castings for Surgical Implants,
- (vi) F 1147-95 "Test Method for Ten-Testing sion of Porous Metal Coatings,
- (vii) F 1378–97 "Specification for
- Shoulder Prosthesis," and (viii) F 1537-94 "Specification for Wrought Cobalt-28 Chromium-6 Molybdenum Alloy for Surgical Implants.

[52 FR 33702, Sept. 4, 1987, as amended at 65 FR 17148, Mar. 31, 2000]

§888.3660 Shoulder joint metal/polysemi-constrained mer cemented prosthesis.

(a) Identification. A shoulder joint metal/polymer semi-constrained cemented prosthesis is a device intended to be implanted to replace a shoulder joint. The device limits translation and rotation in one or more planes via the geometry of its articulating surfaces. It has no linkage across-the-joint. This generic type of device includes prostheses that have a humeral resurfacing component made of alloys, such as cobalt-chromium-molybdenum, and a glenoid resurfacing component made of ultra-high molecular weight polyethylene, and is limited to those prostheses intended for use with bone cement (§ 888.3027).

- (b) *Classification*. Class II. The special controls for this device are:
 - (1) FDA's:
- (i) "Use of International Standard ISO 10993 'Biological Evaluation of Medical Devices—Part I: Evaluation and Testing,"
- (ii) "510(k) Sterility Review Guidance of 2/12/90 (K90-1),"
- (iii) "Guidance Document for Testing Orthopedic Implants with Modified Metallic Surfaces Apposing Bone or Bone Cement."
- (iv) "Guidance Document for the Preparation of Premarket Notification (510(k)) Application for Orthopedic Devices." and
- vices," and
 (v) "Guidance Document for Testing
 Non-articulating, 'Mechanically
 Locked' Modular Implant Components,"
- (2) International Organization for Standardization's (ISO):
- (i) ISO 5832-3:1996 ''Implants for Surgery—Metallic Materials—Part 3: Wrought Titanium 6-aluminum 4-vandium Alloy,''
- (ii) ISO 5832-4:1996 "Implants for Surgery—Metallic Materials—Part 4: Cobalt-chromium-molybdenum casting alloy,"
- (iii) ISO 5832-12:1996 "Implants for Surgery—Metallic Materials—Part 12: Wrought Cobalt-chromium-molybdenum alloy,"
- (iv) ISO 5833:1992 "Implants for Surgery—Acrylic Resin Cements,"
- (v) ISO 5834-2:1998 "Implants for Surgery—Ultra-high Molecular Weight Polyethylene—Part 2: Moulded Forms."
- (vi) ISO 6018:1987 ''Orthopaedic Implants—General Requirements for Marking, Packaging, and Labeling,'' and
- (vii) ISO 9001:1994 ''Quality Systems— Model for Quality Assurance in Design/ Development, Production, Installation, and Servicing,'' and
- (3) American Society for Testing and Materials':
- (i) F 75–92 ''Specification for Cast Cobalt-28 Chromium-6 Molybdenum Alloy for Surgical Implant Material,''

- (ii) F 648-98 "Specification for Ultra-High-Molecular-Weight Polyethylene Powder and Fabricated Form for Surgical Implants,"
- (iii) F 799-96 "Specification for Cobalt-28 Chromium-6 Molybdenum Alloy Forgings for Surgical Implants,"
- (iv) F 1044-95 "Test Method for Shear Testing of Porous Metal Coatings,"
- (v) F 1108-97 "Specification for Titanium-6 Aluminum-4 Vanadium Alloy Castings for Surgical Implants," (vi) F 1147-95 "Test Method for Ten-
- (vi) F 1147–95 "Test Method for Tension Testing of Porous Metal,"
- (vii) F 1378–97 "Standard Specification for Shoulder Prosthesis," and
- (viii) F 1537-94 "Specification for Wrought Cobalt-28 Chromium-6 Molybdenum Alloy for Surgical Implants."

[52 FR 33702, Sept. 4, 1987, as amended at 65 FR 17148, Mar. 31, 2000]

§ 888.3670 Shoulder joint metal/polymer/metal nonconstrained or semiconstrained porous-coated uncemented prosthesis.

(a) Identification. A shoulder joint metal/polymer/metal nonconstrained porous-coated semi-constrained uncemented prosthesis is a device intended to be implanted to replace a shoulder joint. The device limits movement in one or more planes. It has no linkage across-the-joint. This generic type of device includes prostheses that have a humeral component made of alloys such as cobalt-chromium-molybdenum (Co-Cr-Mo) and titanium-aluminum-vanadium (Ti-6Al-4V) alloys, and a glenoid resurfacing component made of ultra-high molecular weight polyethylene, or a combination of an ultra-high articulating molecular weight bearing surface fixed in a metal shell made of alloys such as Co-Cr-Mo and Ti-6Al-4V. The humeral component and glenoid backing have a porous coating made of, in the case of Co-Cr-Mo components, beads of the same alloy or commercially pure titanium powder, and in the case of Ti-6Al-4V components, beads or fibers of commercially pure titanium or Ti-6Al-4V alloy, or commercially pure titanium powder. The porous coating has a volume porosity between 30 and 70 percent, an average pore size between 100 and 1,000 microns, interconnecting porosity, and a porous coating thickness between 500